Research Statement

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Many researchers would be concerned if they learned that some core conclusion of their statistical analysis—such as the sign or statistical significance of some key effect—could be overturned by removing a small fraction, say 0.1%, of their data. Such non-robustness would be particularly concerning if the data were not actually drawn randomly from precisely the population of interest, or if the model may have been misspecified—circumstances that often obtain in the social sciences, for example. Nevertheless, analysts do not routinely check whether ablation of such a small set could overturn their results, in part because the number of possible subsets containing 0.1% of the data points is combinatorially large.

In Giordano [2020] and Broderick et al. [2020], I identify problematic subsets of the data using sensitivity analysis—that is, by forming a linear approximation to how a wide class of statistical estimators depend on their datasets. The key idea is that, although there are a very large number of subsets containing 0.1% of the data points, none of them are very different from the original dataset, and so we expect the linear approximation to work well. I confirm this intuition with finite-sample accuracy bounds in terms of intuitive and verifiable assumptions. I provide an R package [Giordano, 2020] to compute the approximation quickly and automatically using automatic differentiation [Baydin et al., 2017, Maclaurin et al., 2017]. And, with my co-authors, I show that the approximation is capable of detecting meaningful non-robustness in several published econometrics analyses. For example, in a study of microcredit in Mexico [Angelucci et al., 2015], we find that, by removing just 15 households out of 16,561 studied (a change of less than 0.1%), the estimated effect of microcredit changes from negative and statistically insignificant to positive and statistically significant.

In fact, my research shows that many standard, computationally demanding data analysis tasks are also amenable to fast, automatic approximation using sensitivity analysis. For example:

- Cross validation (CV) requires repeatedly leaving out subsets of the observed data and reevaluating a statistical estimator. By forming a Taylor series approximation on the dependence
 of the estimator on the left-out set, I provide fast approximations to CV with finite-sample
 accuracy guarantees [Giordano et al., 2019b].
- Prior specification encodes key assumptions in Bayesian statistics. But Bayesian inference can be sensitive to prior specification, and evaluating the sensitivity of Bayesian posterior expectations to prior specification by re-fitting is typically computationally prohibitive due both to the large space of possible priors (often infinite dimensional), as well as the high computational cost of evaluating even a single posterior approximation. By forming a Taylor series approximation to the dependence of the posterior mean on the prior, I can explore the consequences of alternative prior functional forms at a small fraction of the cost of exact re-fitting [Giordano et al., 2018c, Giordano, 2018].
- Evaluating the frequentist variability of Bayesian estimators formed using Markov Chain Monte Carlo (MCMC) can be particularly important when there is concern about model misspecification. This frequentist variability can be evaluated by the bootstrap, but at the considerable cost of re-running the MCMC chain hundreds of times. By approximating the dependence of the posterior on the data with sensitivity analysis, I compute consistent estimates of the frequentist variance using only a single MCMC chain—orders of magnitude faster than the bootstrap [Giordano and Broderick, 2020b].
- Mean field variational Bayes (MFVB) is a popular posterior approximation method for Bayesian problems which are too large to be tractable by Markov Chain Monte Carlo [Blei et al., 2017, Regier et al., 2019]. However, MFVB approximations provide notoriously poor estimates of posterior uncertainty [Turner and Sahani, 2011]. In Giordano et al. [2018a], I show that accurate posterior covariances can be recovered from MFVB approximations with sensitivity analysis by exploiting a duality between Bayesian covariances and sensitivity.

For the remainder of this essay, I will discuss each of these applications, emphasizing the ways in which I update classical results with intuitive, relevant theory and easy-to-use computational tools.

Robustness to data ablation.

In Giordano [2020] and Broderick et al. [2020], I propose a method to assess the sensitivity of statistical analyses to the removal of a small fraction of the sample. Analyzing all possible data subsets of a certain size is computationally prohibitive, so I provide a finite-sample metric to approximately compute the number (or fraction) of observations that has the greatest influence on a given result when dropped. I provide explicit finite-sample error bounds on my approximation for linear and instrumental variables regressions. At minimal computational cost, the metric provides an exact finite-sample lower bound on sensitivity for any estimator, so any non-robustness my metric finds is conclusive. I demonstrate that non-robustness to data ablation is driven by a low signal-to-noise ratio in the inference problem, is not reflected in standard errors, does not disappear asymptotically, and is not inherently a product of outliers or misspecification.

The approximation works for Z-estimators based on smooth estimating equations, a class which includes ordinary least squares, instrumental variables, generalized method of moments, variational Bayes, and maximum likelihood estimators. Using my R package [Giordano, 2020], the approximation is automatically computable from the specification of the estimating equation alone. By analyzing several published econometric analyses [Angelucci and De Giorgi, 2009, Finkelstein et al., 2012, Meager, 2019], I show that even 2-parameter linear regression analyses of randomized trials can be highly sensitive. While I find some applications are robust, in others the sign of a treatment effect can be changed by dropping less than 1% of the sample even when standard errors are small.

Approximate cross validation.

The error or variability of machine learning algorithms is often assessed by repeatedly re-fitting a model with different weighted versions of the observed data; cross-validation (CV) can be thought of as a particularly popular example of this technique. In Giordano et al. [2019b], I use a linear approximation to the dependence of the fitting procedure on the weights, producing results that can be faster by an order of magnitude than repeated re-fitting. I provide explicit finite-sample error bounds for the approximation in terms of a small number of simple, verifiable assumptions. My results apply whether the weights and data are stochastic or deterministic, and so can be used as a tool for proving the accuracy of the infinitesimal jackknife on a wide variety of problems. As a corollary, I state mild regularity conditions under which the approximation consistently estimates true leave-k-out cross-validation for any fixed k. I demonstrate the accuracy of the approximation on a range of simulated and real datasets, including an unsupervised clustering problem from genomics [Luan and Li, 2003, Shoemaker et al., 2015].

Approximately bootstrapping Bayesian posterior means.

The frequentist (i.e., sampling) variance of Bayesian posterior expectations differs in general from the posterior variance even for large datasets, particularly when the model is misspecified or contains many latent variables [Kleijn and van der Vaart, 2006]. Unlike the posterior variance, the frequentist variance is meaningful even in the presence of misspecification, particularly when the data is known to arise from random sampling [Waddell et al., 2002]. However, the principal existing approach for computing the frequentist variability of MCMC procedures is the bootstrap, which can be extremely computationally intensive due to the need to run hundreds of extra MCMC procedures [Huggins and Miller, 2019].

In Giordano and Broderick [2020a,b], I propose an efficient alternative to bootstrapping an MCMC procedure. My approach is based on the Bayesian analogue of the influence function from the classical frequentist robustness literature. Using results from [Giordano et al., 2018a, 2019b], I show that the influence function for posterior expectations can be easily computed from the posterior samples of a single MCMC procedure and consistently estimates the bootstrap variance. I demonstrate the accuracy and computational benefits of the influence function variance estimates

on array of experiments including an election forecasting model [Gelman and Heidemanns, 2020], the Cormack-Jolly-Seber model from ecology [Kéry and Schaub, 2011], and a large collection of models and datasets from the social sciences [Gelman and Hill, 2006].

Bayesian prior sensitivity.

Prior sensitivity for Markov Chain Monte Carlo. MCMC is arguably the most commonly used computational tool to estimate Bayesian posteriors, which is made still easier by modern black-box MCMC tools such as Stan [Carpenter et al., 2017, Stan Development Team, 2020]. However, a single run of MCMC typically remains time-consuming, and systematically exploring alternative prior parameterizations by re-running MCMC would be computationally prohibitive for all but the simplest models.

My software package, rstansensitivity, [Giordano, 2018, Giordano et al., 2018b], takes advantage of the automatic differentiation capacities of Stan [Carpenter et al., 2015] together with a classical result from Bayesian robustness [Gustafson, 1996, Basu et al., 1996, Giordano et al., 2018a] to provide automatic hyperparameter sensitivity for generic Stan models from only a single MCMC run. I demonstrate the speed and utility of the package in detecting excess prior sensitivity, particularly in a social sciences model taken from Gelman and Hill [2006, Chapter 13.5].

Prior sensitivity for discrete Bayesian nonparametrics. A central question in many probabilistic clustering problems is how many distinct clusters are present in a particular dataset. Discrete Bayesian nonparametric (BNP) mixture models address this question by placing a generative process on cluster assignment, making the number of distinct clusters present amenable to Bayesian inference. However, like all Bayesian approaches, BNP requires the specification of a prior, and this prior may favor a greater or lesser number of distinct clusters.

In Giordano et al. [2018c], I derive prior sensitivity measures for a truncated variational Bayes approximation using ideas from [Gustafson, 1996, Giordano et al., 2018a]. Unlike previous work on local Bayesian sensitivity for BNP [Basu, 2000], I pay special attention to the ability of the sensitivity measures to *extrapolate* to different priors, rather than treating the sensitivity as a measure of robustness *per se*. In work currently in progress [Liu et al., 2020], my co-authors and I apply the approximation from [Giordano et al., 2018c] to an unsupervised clustering problem on a human genome dataset [Huang et al., 2011, Raj et al., 2014], demonstrating that the approximate is accurate, orders of magnitude faster than re-fitting, and capable of detecting meaningful prior sensitivity.

Uncertainty propagation in mean-field variational Bayes.

Mean-field Variational Bayes (MFVB) is an approximate Bayesian posterior inference technique that is increasingly popular due to its fast runtimes on large-scale scientific data sets (e.g., Raj et al. [2014], Kucukelbir et al. [2017], Regier et al. [2019]). However, even when MFVB provides accurate posterior means for certain parameters, it often mis-estimates variances and covariances [Wang and Titterington, 2004, Turner and Sahani, 2011] due to its inability to propagate Bayesian uncertainty between statistical parameters.

In Giordano et al. [2015, 2018a], I derive a simple formula for the effect of infinitesimal model perturbations on MFVB posterior means, thus providing improved covariance estimates and greatly expanding the practical usefulness of MFVB posterior approximations. The estimates for MFVB posterior covariances rely on a result from the classical Bayesian robustness literature that relates derivatives of posterior expectations to posterior covariances and includes the Laplace approximation as a special case. In the experiments, I demonstrate that my methods are simple, general, and fast, providing accurate posterior uncertainty estimates and robustness measures with runtimes that can be an order of magnitude faster than MCMC, including models from ecology [Kéry and Schaub, 2011], the social sciences [Gelman and Hill, 2006], and on a massive internet advertising dataset [Criteo Labs, 2014].

Selected Future work

My research is ideally driven by the needs of my scientific and industry collaborators, and so I expect my future work will be determined to a large part by my colleagues. However, I will now discuss a few directions that I find promising and interesting, and which I believe could be applicable to a diverse set of problems.

The higher-order infinitesimal jackknife for the bootstrap. In the preprint Giordano et al. [2019a], I extend Giordano et al. [2019b] to higher-order Taylor series approximations, providing a family of estimators which I collectively call the higher-order infinitesimal jackknife (HOIJ). In addition to providing higher-quality approximations to CV and extending the results to k-fold CV, the higher-order approach promises to provide a scalable alternative to the bootstrap, a procedure that estimates frequentist variability by repeatedly re-evaluating a model at datasets drawn with replacement from the observed data. The bootstrap is known to enjoy higher-order accuracy in certain circumstances [Hall, 2013], and the HOIJ can approach the bootstrap at a rate faster than the bootstrap approaches the truth. The HOIJ thus promises to make bootstrap inference available to models which are differentiable but too expensive to re-evaluate (e.g. simulation-based models [Baker et al., 2019, Section 2.6]), but also to allow efficient bootstrap-after-bootstrap procedures which that are currently out of reach for all but the simplest statistics [Efron and Tibshirani, 1994].

Scaling sensitivity measures. Sensitivity analysis typically avoids the expense of re-fitting a model, but incurs the expense of solving a large linear system. Thus, extending the benefits of the sensitivity analysis to increasingly large scientific problems requires developing methods to efficiently solve correspondingly large linear systems. Stochastic second-order methods are currently an active research topic in optimization [Agarwal et al., 2017, Berahas et al., 2020], and methods developed therein should apply directly to sensitivity analysis.

Partitioned Bayesian inference. The ideas of [Giordano et al., 2018a] can be naturally extended to approximately propagate uncertainty among separately estimated components of an inference problem. For example, astronomical catalogs are customarily produced with MFVB-like algorithms [Lang et al., 2016, Regier et al., 2019], which take inputs such as the sky background and optical point spread function as fixed inputs, though these quantities are themselves inferred with uncertainty. Viewing all the separate inference procedures as a sequential quasi-MFVB objective, one could directly apply the techniques of [Giordano et al., 2018a] to propagate the uncertainty from the modeling inputs to the astronomical catalog's uncertainty.

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